Psychological Burden amongst Caregivers of Patients on Maintenance Haemodialysis

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ABSTRACT

Long-term caregiving is a difficult job, risking mental health of involved individuals. This study was planned to document magnitude of the problem at Pak Emirates Military Hospital, Rawalpindi, with an aim to improve quality of life in caregivers and patients alike. From January to June 2019, caregivers accompanying haemodialysis patients were selected using consecutive sampling technique. Those unwilling were excluded. Psychological burden was assessed using Zarit Burden Interview (ZBI) questionnaire. There were 98 caregivers, including 71 (72.45%) males and 27 (27.55%) females, aged 33.13 ± 9.98 years. There was little/no burden in 27 (27.55%) caregivers; 52 (53.06%) had mild to moderate, 18 (18.37%) had moderate to severe and 3 (3.06%) had severe psychological burden. Mean ZBI score was 30.00 ± 12.57. Males had lower scores than females (28.01 ± 11.18 vs. 35.22 ± 14.65, respectively; p = 0.026). Gender of caregivers significantly predicted ZBI scores; while their age, job status, education, monthly income and duration of caregiving did not.

Key Words: Caregivers, Chronic kidney disease, Haemodialysis, Long-term care, Psychological burden.

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The burden of end-stage renal disease (ESRD) is increasing progressively, proportionate to the growing prevalence of major risk factors including diabetes and hypertension.¹ Renal transplant is the treatment of choice, known to improve both morbidity and mortality. Many patients cannot afford this because of financial constraints or shortage of donors. This leaves them with the only option of long-term haemodialysis (HD). They have to attend in-centre HD sessions three times a week for an indefinite period, a difficult task. Caregivers play an important role in looking after patients with ESRD. This could include taking care of their needs at home as well as accompanying them to HD centre. These roles assume greater importance in Pakistan because of cultural values. While managing patients, we often ignore their family members. Another important aspect is the perception of caregivers’ contributions by the patients themselves. This is significant as this could affect the quality of life of the patients themselves. Whereas, we now have a lot of local data assessing psychiatric morbidity amongst patients with ESRD, not many studies from Pakistan have looked into the problems faced by their caregivers.

We, therefore, planned this study to determine the magnitude of the problem in our setup. The results would help create more awareness and open avenues for treatment of stressed caregivers. Such an action would indirectly improve outcomes for ESRD patients as well.

This cross-sectional study was carried out at the Department of Nephrology, Pak Emirates Military Hospital, Rawalpindi, from January to June 2019 after obtaining approval from the local Ethics Review Committee (Reference number A/28/PEMH/EC-19/21). We enlisted all patients who had been on maintenance HD at our centre for at least three months and were accompanied by caregivers at the HD unit. We invited their caregivers (only family members living with the patients) for inclusion in this study, using consecutive sampling technique.

Only one caregiver for any given patient on HD was approached. Caregivers not providing consent, those known to have anxiety, depression or any other psychiatric illness, domestic servants and those helping the patients only with transportation to and from HD unit, were excluded. A minimum sample size of 97 was calculated with Free Statistics Calculators version 4.0, using an anticipated effect size (f²) of 0.15, power of 0.8, alpha of 0.05 and 6 predictors. We collected demographic data on six factors that could potentially predict psychological burden in the caregivers: their age, gender, occupation, level of education, monthly income and the time period they had been coming to the hospital as caregivers.

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Psychological burden was assessed using ZBI. This questionnaire contains 22 items, each having five responses scored from 0 to 4. Total scores up to 21 indicate little or no burden, 21-40 indicate mild to moderate burden, 41-60 indicate moderate to severe burden, whereas 61-88 reflect severe burden.

The questionnaire was translated to Urdu and then back-translated to English by two separate doctors. The original and back-translated version were compared by the first author to ensure that no significant difference had emerged during this process. We then used the Urdu version for this study. Patients literate enough to comprehend the questionnaire self-reported. Others were asked all the questions in direct face-to-face interviews by the second author.

Data was analysed using IBM SPSS Statistics for Windows, Version 20.0 (IBM Corp, Armonk, NY). Quantitative data with parametric distribution was described as mean ± standard deviation (SD), whereas non-parametric data was described as median and range. Standard multiple regression was carried out to determine the relationship between ZBI scores and the six possible predictors. P values ≤0.05 were considered statistically significant.

The mean age of 98 caregivers was 33.13 ± 9.98 years. There were 71 (72.45%) males and 27 (27.55%) females. Most (n=66; 67.35%) were working, while 32 (32.65%) did not have any job. Mean education level was 10.88 ± 3.72 years. Median monthly income was Rs. 18,000 (range: 0-180,000). Mean attendance vintage was 12 months (range: 0-120 months). Majority of the attendants (n=90; 91.84%) accompanied patients on more than 50% of the sessions, whereas the rest (8; 8.16%) were present during less than 50% sessions. There was little/no psychological burden in 27 (27.55%) caregivers; 52 (53.06%) had mild to moderate, 18 (18.37%) had moderate to severe and 3 (3.06%) had severe psychological burden. Mean ZBI score was 30.00 ± 12.57. Mean scores for individual items of this scale are shown in Table I. Males had lower scores than females (28.01 ± 11.18 vs. 35.22 ± 14.65, respectively; p= 0.026). As shown in Table II, gender of the caregivers was the only factor significantly predicting total ZBI scores.

Caregivers are almost universally required for ESRD patients considering their advanced age, presence of multiple co-morbidities and intricacies of HD. Looking after these patients is an overwhelming experience, with greater expectations keeping in view the traditional Eastern cultural values. Close family members of HD patients are thus an integral part of the management team, sharing important responsibilities in care. This study has highlighted psychological burden in a vast majority (72.45%) of caregivers. Most of them perceived only mild to moderate burden. Our results are comparable to figures reported in a study done at other dialysis setups in Rawalpindi five years ago. Caregiving burden was noted in 80% of those cases, mostly mild to moderate.
Left unaddressed, this burden would eventually translate into poor quality of life.

Majority of the caregivers were males. The male preponderance in our study population could be because of the physical strength required to move patients between home and dialysis centre. It is very much possible that the roles get changed at homes, where females could look after other needs of these patients. We do not have further information on this for our study population. Female caregivers had higher ZBI scores in this study, similar to results of a Turkish study. The most plausible explanation would be the stress of looking after patients in addition to the responsibility of carrying out household chores and taking care of other family members. Strangely, none of the other parameters studied was associated with psychological burden. The results are in contrast to generally available literature. Farzi et al. documented greater burden in working caregivers from Iran. It is very much possible that the caregivers in our cohort had adopted coping strategies to minimise the impact of work on their daily lives. Caregivers’ income also did not have any relationship with psychological burden in this study. Similar results were reported by Hoang et al. in a study done from Vietnam; whereas, Mashayekhi et al. described a significant relationship of income with caregiver burden score amongst Iranian patients.

ZBI has been used to assess caregiver burden in a number of clinical conditions. Amongst caregivers of patients with heart failure, it has been shown to have a good internal consistency (Cronbach's α= 0.921) with good item-total (r = 0.395-0.764) and item-item (mean r = 0.365) correlations.

Apart from the inherent problems associated with a cross-sectional study design, this study is limited by the absence of a control group comprising of healthy individuals. It is generally believed that the total time spent caring for patients in a day is a predictor of psychological burden. We focused only on caregivers’ attendance at HD sessions and did not record data about relevant activities while at home. Moreover, we do not have data to suggest the extent to which the patients could care for themselves. Data on transportation used by patients is also missing. Some of the caregivers had to be interviewed as they could not self-administer the questionnaire. This could have produced information bias.

In short, caregivers’ psychological burden is a significant problem in haemodialysis patient populations, more so amongst females. This necessitates periodic screening, with an aim to help affected persons in reducing the mental stress.

**ETHICAL APPROVAL:**
Ethical approval was obtained from Ethical Committee of Pak Emirates Military Hospital, Rawalpindi, vide approval letter No. A/28/PEMH/EC-19/21.

**PATIENTS’ CONSENT:**
Informed written consents were obtained from all study participants.

**CONFLICT OF INTEREST:**
The authors declared no conflict of interest.

**AUTHORS’ CONTRIBUTION:**
ARA: Interpreted data, critically revised the manuscript, approved the final version, agreed to be accountable for all aspects.
TT: Conceived the study, analysed data, drafted the manuscript, approved the final version, agreed to be accountable for all aspects.
AWM: Designed the study, acquired data, drafted the manuscript, approved the final version, agreed to be accountable for all aspects.
SU: Analysed data, drafted the manuscript, approved the final version, agreed to be accountable for all aspects.

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